

Interleukin-5 (IL-5) Inhibitors Prior Authorization with Quantity Limit Program Summary

This program applies to FlexRx Closed, FlexRx Open, FocusRx, GenRx Closed, GenRx Open, Health Insurance Marketplace, and KeyRx formularies.

This is a FlexRx Standard and GenRx Standard program.

The BCBS MN Step Therapy Supplement also applies to this program for all Commercial/HIM lines of business.

POLICY REVIEW CYCLE

 Effective Date
 Date of Origin

 05-01-2024
 07-01-2019

FDA APPROVED INDICATIONS AND DOSAGE

Agent(s)	FDA Indication(s)	Notes	Ref#
Fasenra®	Add-on maintenance treatment of patients with severe asthma aged 12 years and older, and with an eosinophilic phenotype		2
(benralizumab)	Limitations of use:		
Injection for subcutaneous use	 Treatment of other eosinophilic conditions Relief of acute bronchospasm or status asthmaticus 		
Nucala®	Add-on maintenance treatment of patients aged 6 years and older with severe asthma and with an eosinophilic phenotype		1
(mepolizumab) Injection for	Treatment of adult patients with eosinophilic granulomatosis with polyangiitis (EGPA)		
subcutaneous use	Treatment of adult and pediatric patients aged 12 years and older with hypereosinophilic syndrome (HES) for greater than or equal to 6 months without an identifiable non-hematologic secondary cause		
	Add-on maintenance treatment of chronic rhinosinusitis with nasal polyps (CRSwNP) in adult patients 18 years of age and older with inadequate response to nasal corticosteroids		
	Limitation of use:		
	 Not for relief of acute bronchospasm or status asthmaticus 		

See package insert for FDA prescribing information: <u>https://dailymed.nlm.nih.gov/dailymed/index.cfm</u>

CLINICAL RATIONALE

Asthma	Asthma is a chronic inflammatory disorder of the airways.(3,5) It is characterized by variable and recurring clinical symptoms, airflow obstruction, bronchial
	hyperresponsiveness, and underlying inflammation.(3) Symptoms of asthma include wheezing, coughing, recurrent difficulty breathing, shortness of breath, and chest

tightness. Generally, these symptoms will occur or worsen with exposure to allergens and irritants, infections, exercise, changes in weather, stress, or menstrual cycles. Guidelines recommend the use of detailed medical history, physical examination, and spirometry to make a diagnosis of asthma.(3,5)
The Global Initiative for Asthma (GINA) guidelines recommend a stepwise approach for managing asthma. Long-term goals for asthma management are to achieve good control of symptoms, maintain normal activity level, and to minimize the future risk of exacerbations, fixed airflow limitation, and side-effects.(5) IgE is the antibody responsible for activation of allergic reactions and is important to the pathogenesis of allergic asthma and the development and persistence of inflammation. GINA guidelines define moderate asthma as that which is well controlled with Step 3 or Step 4 treatment (e.g., low- or medium-dose inhaled corticosteroids [ICS] in combination with a long-acting beta agonist [LABA] in either treatment track). Severe asthma is defined as asthma that remains uncontrolled despite optimized treatment with high- dose ICS-LABA, or that requires high-dose ICS-LABA to prevent it from becoming uncontrolled. Severe asthma must be distinguished from asthma that is difficult to treat due to inadequate or inappropriate treatment, or persistent problems with adherence or comorbidities such as chronic rhinosinusitis or obesity, as they need very different treatment compared with if asthma is relatively refractory to high-dose ICS- LABA or even oral corticosteroids (OCS). Early initiation of low dose ICS in patients with asthma has led to greater improvement in lung function than initiation of ICS after symptoms have been present for more than 2 to 4 years. The 2023 GINA guidelines recommend every adult and adolescent with asthma should receive ICS- containing controller medication to reduce the risk of serious exacerbation, even in patients with infrequent symptoms.(5)
2023 GINA STEP recommendations for adults and adolescents (12 years of age and over) are intended to reduce the risk of serious exacerbations and are broken into two tracks based on reliever therapy.
Track 1 is the preferred approach recommended by GINA, because using low dose ICS-formoterol as reliever reduces the risk of exacerbations compared with regimens with short-acting β 2-agonist (SABA) as reliever, and is a simpler regimen. Note ICS-formoterol should not be used as the reliever by patients taking any other (non-formoterol) ICS-LABA or ICS-LAMA:(5)
• Step 1:
• As-needed low dose ICS-formoterol
 Step 2. As-needed low dose ICS-formoterol
 Step 3: address and treat modifiable risk factors (e.g., adherence, technique) before considering step up
Maintenance: low dose ICS-formoterol Believer: as-needed low dose ICS-formoterol
Step 4:
 Maintenance: medium dose ICS-formoterol Reliever: as-needed low dose ICS-formoterol
 Step 5: patients with uncontrolled symptoms and/or exacerbations despite Step 4 treatment should be assessed for contributory factors, have their
treatment optimized, and be referred for expert assessment including severe
 Maintenance: consider high dose ICS-formoterol
 Reliever: as-needed low dose ICS-formoterol Add-on LAMA for patients greater than or equal to 18 years (greater
than or equal to 6 years for tiotropium) in separate or combination inhalers
 Refer for phenotypic assessment +/- biologic therapy Add on anti Infe for source allegate attempts
 Add-on anti-igE for severe allergic astima SC omalizumab in patients greater than or equal to 6
years

 Add-on anti-interleukin (L)S or anti-LSR or anti-IL4R for severe eosinophilic/Type 2 asthma Anti-LS: SC menolizumab for patients greater than equal to 5 years OR. Yreslizumab for patients greater than or equal to 13 years Anti-LRS: SC benarizumab for patients greater than equal to 6 years Anti-LRS: SC benarizumab for patients greater than equal to 6 years Add-on anti-thymic stromal lymphopoletin (TSLP) for sever asthma St pays anti-transmission of the strong lymphopoletin (TSLP) for sever asthma St pays Add-on anti-thymic stromal lymphopoletin (TSLP) for sever asthma St pays anti-transmission of the strong lymphopoletin (TSLP) for sever asthma St pays anti-transmission of the strong lymphopoletin (TSLP) for sever asthma St pays anti-transmission of the strong lymphopoletin (TSLP) for sever asthma St pays anti-transmission of the strong lymphopoletin (TSLP) for sever asthma St pays anti-transmission of the strong lymphopoletin (TSLP) for sever asthma St pays anti-transmission of the strong lymphopoletin (TSLP) for sever asthma Take Z is an alternative approach if Track 1 is not possible or is not preferred by patient with the concluster should consider whether the patient is likely to be adherent with their controller therapy; if not, they will be exposed to the higher ris exacerbations with SABA-only treatment:(5) Step 1: Take ICS whenever SABA taken Reliever: as-needed ICS-SABA or as needed SABA Alternative options with limited indications, or less evidence for efficacy and/or safety: Low dose ICS-Whenever SABA taken Daily low-dose ICS-LABA as initial therapy leads to faster im		
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Step 4: O Preferred maintenance: medium/high dose ICS-LABA O Preferred reliever: as-needed ICS-SABA or as-needed SABA		 Low-dose ICS plus LTRA but review US FDA boxed warning For adults with rhinitis who are allergic to house dust mite and have FEV1 > 70% predicted, consider adding SLIT
		Step 4: • Preferred maintenance: medium/high dose ICS-LABA • Preferred reliever: as-needed ICS-SABA or as-needed SABA
 Alternative options: Add-on LAMA for patients greater than or equal to 18 years (greater than or equal to 6 years for tiotropium my mist inhaler) 		 Alternative options: Add-on LAMA for patients greater than or equal to 18 years (greater than or equal to 6 years for tiotropium my mist inhaler)

 Before considering add-on LAMA for patients with exacerbations, increase ICS dose to at least medium For adults with rhinitis who are allergic to house dust mite and have FEV1 > 70% predicted, consider adding sublingual immunotherapy (SLIT)
 Step 5: patients with uncontrolled symptoms and/or exacerbations despite Step 4 treatment should be assessed for contributory factors, have their treatment optimized, and be referred for expert assessment including severe asthma phenotype, and potential add on treatment Maintenance: medium/high dose ICS-LABA
 Reliever: as-needed ICS-SABA or as-needed SABA Add-on LAMA for patients greater than or equal to 18 years (greater than or equal to 6 years for tiotropium) in separate or combination inhalers
 Refer for phenotypic assessment +/- biologic therapy Add-on anti-IgE for severe allergic asthma SC omalizumab in patients greater than or equal to 6
years Add-on anti-interleukin (IL)5 or anti-IL5R or anti-IL4R for severe eosinophilic/Type 2 asthma
Anti-ILS: SC mepolizumab for patients greater than or equal to 6 years OR IV reslizumab for patients greater than or equal to 18 years of age
 Anti-ILSR: SC bernalization for patients greater than or equal to 12 years Anti-IL4R: SC dupilumab for patients greater than or equal to 6 years
 Add-on anti-thymic stromal lymphopoietin (TSLP) for severe asthma SC tozonolumab for patients greater than or equal to a severe asthmatic stromatic st
 Or Sector period and for patients greater than or equal to 12 years Add-on azithromycin three days/week reduces exacerbations, but increases antibiotic resistance
 Maintenance OCS should only be used as last resort, because short- term and long-term systemic side-effects are common and serious
2023 GINA STEP recommendations for children (6 to 11 years of age) are intended to reduce the risk of serious exacerbations:(5)
 Step 1: Low dose ICS taken whenever SABA taken Reliever: as needed SABA
 Step 2: Preferred: daily low dose ICS Preferred reliever: as needed SABA Alternative options: Low-dose ICS whenever SABA is taken using separate inhalers Daily I TRA are less effective for exacerbation reduction. Advise
 parents about US FDA warning on montelukast Step 3: after checking inhaler technique and adherence, and treating modifiable risk factors (any of the following):
 Medium-dose ICS maintenance plus as-needed SABA Low-dose ICS-LABA maintenance plus as-needed SABA Maintenance and reliever therapy (MART) with a very low dose of budesonide-formoterol DPI
 Step 4: Individual children's responses vary, so each of the Step 3 options may be tried before considering a step-up to Step 4. Refer for expert advice Preferred: medium dose ICS-LABA plus as-needed SABA Preferred: low dose ICS-formoterol MART plus as-needed low-dose ICS-formoterol Alternative options:
 Add-on tiotropium

Add-on LTRA
 Refer for phenotypic assessment with or without higher dose ICS-LABA Reliever: as needed SABA (or ICS-formoterol reliever for MART) Add on therapy with anti-IgE or anti-IL4R, anti-IL5 As a last resort consider add on low dose OCS but consider side effects
Severe Asthma Phenotype and Eosinophilic Asthma Subphenotype
Roughly 3% to 10% of adults with asthma have severe asthma as defined by the GINA 2023 guidelines.(5) The European Respiratory Society (ERS)/American Thoracic Society (ATS) guidelines (2014; updated 2020) and the National Asthma Education and Prevention Program Coordinating Committee Expert Panel Working Group mirror the GINA definition of severe asthma, and defined uncontrolled asthma for adult and pediatric patients 5 years of age and over:(3,19)
 Frequent severe exacerbations (i.e., two or more bursts of systemic corticosteroids within the past 12 months) Serious exacerbations (i.e., at least one hospitalization, intensive care unit
 stay, or mechanical ventilation in the past 12 months) Airflow limitation (i.e., FEV1 less than 80% predicted)
 Asthma that worsens upon tapering of high-dose ICS or systemic corticosteroids
A specialist, preferably in a multidisciplinary severe asthma clinic (if available) performs further assessment, which includes the patient's inflammatory phenotype (i.e., Type 2 or non-Type 2).(5)
Type 2 inflammation is characterized by the presence of cytokines such as interleukin (IL)-4, IL-5, and IL-13, which are often produced by the adaptive immune system on recognition of allergens. It is also characterized by eosinophilia or increased fraction of exhaled nitric oxide (FeNO) and may be accompanied by atopy. In many patients with asthma, Type 2 inflammation rapidly improves when ICS are taken regularly and correctly; this is classified as mild or moderate asthma. In severe asthma, Type 2 inflammation is considered refractory if any of the following are found while the patient is taking high dose ICS or daily OCS:(5)
Blood eosinophils greater than or equal to 150 cells/microliter
 FeNO greater than or equal to 20 ppb Sputum eosinophils greater than or equal to 2% Asthma is clinically allergen-driven
Biologic agents should be considered as add-on therapy for patients with refractory Type 2 inflammation with exacerbations or poor symptom control despite taking at least high dose ICS/LABA, and who have allergic or eosinophilic biomarkers or need maintenance OCS.(5) 2023 GINA recommends the biologics below based on patient eligibility factors:
 Anti-IgE (omalizumab): Sensitization on skin prick testing or specific IgE Total serum IgE and weight within dosage range Exacerbations in the last year Anti-IL5/Anti-IL5R (benralizumab, mepolizumab, reslizumab): Exacerbations in the last year Blood eosinophils greater than or equal to 150 cells/microliter (for benralizumab and mepolizumab) or greater than or equal to 300 cells/microliter (for reslizumab) Anti-IL4R (dupilumab): Exacerbations in the last year

	 Blood eosinophil greater than or equal to 150 cells/microliter but less than or equal to 1500 cells/microliter, or FeNO greater than or equal to 25 ppb, or taking maintenance OCS Anti-TSLP (tezepelumab): Exacerbations in the last year
	Patient response should be evaluated 4 months after initiating therapy and follow up should occur every 3 to 6 months thereafter. 2023 GINA recommends the following step-down therapy process in patients responding well to targeted biologic therapy:(5)
	 Reevaluate the need for each asthma medication every 3 to 6 months, but inhaled therapy should not be completely stopped Oral treatments: gradually decreased starting with OCS due to significant adverse effects. Inhaled treatments: consider reducing ICS dose after 3 to 6 months, but do not completely stop inhaled therapy. Continue at least medium dose ICS and remind patients of the importance of continued inhaled controller therapy Biologic treatments: trial withdrawal after 12 months of treatment and only if patient's asthma remains well controlled on medium dose ICS, and for allergic asthma, there is no further exposure to a previous allergic trigger
Eosinophilic Granulomatosis with Polyangiitis (EGPA)	Eosinophilic granulomatosis with polyangiitis (EGPA), formally known as Churg-Strauss Syndrome, is a rare small-vessel vasculitis that occurs in patients with asthma and eosinophilia and is histologically characterized by tissue eosinophilia, necrotizing vasculitis and eosinophil-rich granulomatous inflammation. EGPA is an anti-neutrophil cytoplasmic antibody (ANCA)-associated vasculitis, characterized by asthma, eosinophilia and granulomatous or vasculitic involvement of several organs. Current practice relies on recommendations and guidelines addressing the management of ANCA-associated vasculitis and not specifically developed for EGPA.(20) The main clinical features of EGPA are late-onset allergic rhinitis and asthma, increased blood eosinophil count, and vasculitis manifestations, some of which can be life threatening. Once EGPA is suspected based on clinical findings of asthma with eosinophilia, asthma with systemic manifestations, or even eosinophilia with extrapulmonary disease, a biopsy demonstrating small or medium sized vessel vasculitis strongly supports the diagnosis of EGPA. Skin, nerve, and muscle are among the most common biopsied tissues, but endomyocardial, renal, and gastrointestinal biopsies may also be useful. Antineutrophil cytoplasm antibody (ANCA) testing is also recommended. ANCA positivity is highly suggestive of EGPA, but ANCA negative results do not rule out its diagnosis.(6)
	The clinical phenotype of EGPA is quite heterogeneous and the diagnosis is not always straightforward. Anti-neutrophil cytoplasmic antibodies (ANCA), usually against myeloperoxidase (MPO), are detectable in approximately 40% of the cases and are associated with a different frequency of clinical manifestations: features of vasculitis, particularly glomerulonephritis, peripheral neuropathy and purpura, occur more often in ANCA-positive patients, whereas the so-called eosinophilic features such as cardiac involvement and gastroenteritis are more frequent in ANCA-negative patients.(20)
	There are two types of classifications used for the diagnosis of EGPA. The first and most commonly used classification is by the American College of Rheumatology (ACR). ACR has established six criteria for the classification of EGPA in a patient with documented vasculitis. The presence of four or more of these criteria can establish a diagnosis of EGPA:(7)
	 Asthma (a history of wheezing or diffuse high-pitched rales on expiration) Eosinophilia (greater than 10% eosinophils on white blood cell differential count) Mononeuropathy (including multiplex), multiple mononeuropathies, or polyneuropathy attributed to a systemic vasculitis Migratory or transient pulmonary infiltrates detected radiographically

•	Paranasal	sinus	abnormality
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• Biopsy containing a blood vessel showing the accumulation of eosinophils in extravascular areas

The Lanham criteria is also used for the diagnosis of EGPA. The Lanham criteria requires the patient to have all three of the following: asthma, peak peripheral blood eosinophilia in excess of 1500 cells/microliter, and systemic vasculitis involving two or more extra-pulmonary organs.(7,8)

The American College of Rheumatology/European Alliance of Associations for Rheumatology developed classification criteria for EGPA broken into clinical criteria as well as laboratory and biopsy criteria. Considerations when applying these criteria(20)

- These classification criteria should be applied to classify a patient as having EGPA when a diagnosis of small- or medium-vessel vasculitis has been made
- Alternate diagnoses mimicking vasculitis should be excluded prior to applying the criteria

	points
Clinical Criteria	
Obstructive airway disease	+3
Nasal Polyps	+3
Mononeuritis multiplex	+1
Laboratory and Biopsy Criteria	
Blood eosinophil count greater than or equal to 1 X 10^9/liter	+5
Extravascular eosinophilic-predominant inflammation on biopsy	+2
Positive test for cytoplasmic antineutrophil cytoplasmic antibodies (cANCA)	-3
Hematuria	-1

Sum the scores for the 7 items, if present. A score of greater than or equal to 6 is needed for classification of EGPA

The Five-Factor Score (FFS) predicts the risk of mortality in patients with an established diagnosis of EGPA, as well as polyarteritis nodosa microscopic polyangiitis or GPA. It includes five factors associated with shortened overall survival, namely, renal insufficiency (serum creatinine > 1.58 mg/dl), proteinuria > 1 g per day, cardiomyopathy, gastrointestinal involvement and central nervous system (CNS) involvement. The FFS considers clinical manifestations only at the time of diagnosis, the appearance of new manifestations during follow-up should also be taken into account when choosing remission-induction regimens for disease flares. New-onset active EGPA is considered severe if FFS is greater than or equal to 1 or there is presence of peripheral neuropathy, alveolar hemorrhage or other organ-or life-threatening manifestations. For relapsing EGPA severe disease consists of severe systemic relapse and non-severe is respiratory relapse alone or non-severe systemic relapse.(20)

For remission induction in patients with new-onset, active EGPA, glucocorticoids should be administered as initial therapy. In patients with severe disease cyclophosphamide or rituximab and/or disease modifying anti-rheumatic drugs (DMARDs) should be added to glucocorticoid therapy. Remission maintenance for non-severe disease guidelines recommend glucocorticoids plus mepolizumab. Remission maintenance for severe disease guidelines recommend glucocorticoids plus rituximab and/or mepolizumab and or DMARDs. Although the evidence supporting the use of traditional

	immunosuppressants for remission maintenance in non-severe EGPAS is scarce, these agents are often used in routine clinical practice.(20)
	Treatment for relapsing EGPA in non-severe disease glucocorticoids alone or glucocorticoids plus mepolizumab along with optimization of inhaled therapies. Treatment of relapsing severe disease high-dose oral glucocorticoids plus cyclophosphamide or rituximab is recommended.(20)
	Refractory EGPA is defined as unchanged or increased disease activity after 4 weeks of appropriate remission-induction therapy. The persistence or worsening of systemic manifestations should be distinguished from that of respiratory manifestations. Mepolizumab in combination with glucocorticoids is recommended to induce remission in patients with relapsing-refractory EGPA without organ-or life-threatening manifestations. Mepolizumab can also be used for remission maintenance, particularly in patients requiring a daily prednisone greater than or equal to 7.5 mg for the control of their respiratory manifestations.(20)
Hypereosinophilic Syndrome (HES)	The eosinophilias encompass a broad range of non-hematologic (secondary or reactive) and hematologic (primary or clonal) disorders with potential for end-organ damage. Hypereosinophilia (HE) has generally been defined as peripheral blood eosinophil count greater than 1500 cells/microliter, OR pathologic confirmation of tissue HE by at least one of the following: percentage of eosinophils in bone marrow section exceeds 20% of all nucleated cells, marked deposition of eosinophil granule proteins is found, or tissue infiltration by eosinophils is extensive in the opinion of the pathologist.(12) To establish a diagnosis of HES, all three of the following criteria must be met:(11,12,13)
	 Criteria for HE fulfilled Evidence of HE-related organ damage (e.g., fibrosis of lung, heart, digestive tract, skin, etc; thrombosis with or without thromboembolism; cutaneous erythema, edema/angioedema, ulceration, pruritis, or eczema; peripheral or central neuropathy with chronic or recurrent neurologic deficit; other organ system involvement such as liver, pancreas, kidney) Exclusion of secondary (non-hematologic) causes of eosinophilia (e.g., infection, allergy/atopy, medications, collagen vascular disease, metabolic [e.g., adrenal insufficiency], solid tumor/lymphoma)
	Although the clinical manifestations can be similar irrespective of the cause of the eosinophilia, the choice of the initial therapeutic agent(s) for a given patient depends mainly on whether the patient has clinical features consistent with a myeloid disorder. Patients with myeloid variants of HES (e.g., PDGFRA-positive HES) often have an aggressive course with disabling complications and high mortality in the absence of treatment, and are treated initially with imatinib; those with other types of HES are treated with an initial trial of glucocorticoids.(11,12,13,14) Oral corticosteroids have been used for decades in the treatment of HES and, with the exception of imatinib for PDGFRA-associated HES as noted above, remain the first-line treatment for most patients. Hydroxyurea is a typical second-line agent, whether used as monotherapy or in conjunction with corticosteroids. Additional immunomodulatory and cytotoxic agent options include interferon-a, azathioprine, cyclosporine, methotrexate, and tacrolimus.(12,13,14)
	Despite the wide variety of commercially available immunomodulatory and cytotoxic agents, a significant proportion of patients with HES are treatment-refractory or experience treatment-related toxicity. Monoclonal anti–IL-5 antibody therapy for HES has a number of unique advantages related to the specificity of IL-5 for the eosinophil lineage.(12,13,14)
Chronic Rhinosinusitis with Nasal Polyposis	Chronic rhinosinusitis with nasal polyposis (CRSwNP) is an inflammatory condition affecting the paranasal sinuses. The International Consensus Statement on allergy and rhinology: Rhinosinusitis indicates that the diagnostic criteria for chronic rhinosinusitis (CRS) consist of ALL the following:(18)

	Symptoms greater than or equal to 12 weeks
	Two of the following symptoms:
	 Nasal discharge (rninorrhea or post-hasal drainage) Nasal obstruction or congestion
	 Hyposmia (loss or decreased sense of smell)
	• Facial pressure or pain
	One or more of the following findings: Description of the following findings:
	 Evidence of inflammation on hasal endoscopy or computed tomography
	 Evidence of purulence coming from paranasal sinuses or ostiomeatal
	complex
	Sinus computed tomography (CT) and/or pasal endoscopy are needed to determine
	the presence of sinonasal inflammation and nasal polyps. The exact cause of CRSwNP
	is unknown, but biopsies of nasal polyps have shown elevated levels of
	eosinophils.(15)
	First line therapy for CRSwNP consists of pasal saline irrigation in combination with
	intranasal corticosteroids.(15,16,17) The American Academy of Family Physicians
	notes that no one intranasal corticosteroid is superior to another or that increased
	dosing provides greater effectiveness. The American Academy of Otolaryngology
	intranasal corticosteroids after 3-months of appropriate use.(17) Short courses of oral
	corticosteroids (up to three weeks) can improve sinonasal symptoms and endoscopic
	findings. Surgical intervention may be required in patients in which medical therapy is
Efficacy	Acthmo
Lincacy	Astima
	Fasenra
	Benralizumab was approved through 3 confirmatory clinical trials. Trial 1 and Trial 2
	were exacerbation trials in patients 12 years of age and older. All subjects continued
	their background asthma therapy throughout the duration of the trials. The primary
	ICS and LABA. Asthma exacerbation was defined as a worsening of asthma requiring
	use of oral/systemic corticosteroids for at least 3 days, and/or emergency department
	visits requiring use of oral/systemic corticosteroids and/or hospitalization. For patients
	on maintenance oral corticosteroids, an asthma exacerbation requiring oral
	corticosteroids for at least 3 days or a single depo-injectable dose of corticosteroids. In
	Trial 1, 35% of patients receiving benralizumab experienced an asthma exacerbation
	compared to 51% on placebo. In Trial 2, 40% of patients receiving benralizumab
	experienced an astrima exacerbation compared to 51% on placebo.(2)
	Trial 3 was a randomized OCS reduction trial in asthma patients. Patients were
	required to be treated with daily OCS (7.5 to 40 mg per day) in addition to regular use
	of high-dose ICS and LABA with or without additional controller(s). The trial included
	an 8-week run-in period during which the OCS was titrated to the minimum effective
	asthma control was assessed by the investigator based on a patient's FEV1, peak
	expiratory flow, nighttime awakenings, short-acting bronchodilator rescue medication
	use or any other symptoms that would require an increase in OCS dose. Fasenra
	achieved greater reductions in daily maintenance OCS dose while maintaining asthma control compared to placebo (median reduction of 75% for Fasenra vs 25% for
	placebo).(2)
	Nucala
	The efficacy of menolizumab for the treatment of severe eosinophilic asthma was
	established in three double-blind, randomized, placebo-controlled trials: A dose- ranging and exacerbation reduction trial (trial 1) and two confirmatory trials (trial 2

and 3). All subjects continued their background asthma therapy throughout the duration of the trials. Trial 1 enrolled subjects with uncontrolled asthma despite use of high dose inhaled corticosteroids (ICS) plus additional controller(s), with or without OCS. Trial 2 was a placebo- and active-controlled trial in subjects with asthma not adequately controlled on high-dose inhaled corticosteroids plus additional controller(s) with or without OCS. The primary end point for trial 1 and 2 was frequency of asthma exacerbations. Compared to placebo, subjects receiving mepolizumab experienced significantly fewer exacerbations and had a longer time to first exacerbation.(1)

Trial 3 was an OCS-reduction study in asthma patients who required daily OCS in addition to regular controller medications. The primary end point was percent reduction of OCS dose during weeks 20 to 24 without loss of asthma control. The baseline mean oral corticosteroid use was similar between the Nucala and placebo group. Overall, mepolizumab achieved greater reduction in oral corticosteroid use while maintaining asthma control when compared to placebo. However, the difference between the mepolizumab and placebo groups was not statistically significant.(1)

EGPA

Nucala

A total of 136 subjects with EGPA were evaluated in a randomized, placebo-controlled, multicenter, 52-week trial. Subjects enrolled had a diagnosis of EGPA for at least 6 months prior to enrollment with a history of relapsing or refractory disease and were on a stable dosage of oral prednisolone or prednisone of greater than or equal to 7.5 mg/day (but not greater than 50 mg/day) for at least 4 weeks prior to enrollment. Subjects received 300 mg of mepolizumab or placebo administered subcutaneously once every 4 weeks while continuing their stable OCS therapy. Starting at Week 4, OCS was tapered during the treatment period at the discretion of the investigator. The co-primary endpoints were the total accrued duration of remission over the 52-week treatment period, defined as Birmingham Vasculitis Activity Score (BVAS) = 0 (no active vasculitis) plus prednisolone or prednisone dose less than or equal to 4 mg/day, and the proportion of subjects in remission at both Week 36 and Week 48 of treatment. The BVAS is a clinician-completed tool to assess clinically active vasculitis that would likely require treatment, after exclusion of other causes.(1)

A significantly higher proportion of subjects receiving mepolizumab achieved remission at both Week 36 and Week 48 compared with placebo. In addition, significantly more subjects receiving mepolizumab achieved remission within the first 24 weeks and remained in remission for the remainder of the 52-week study treatment period compared with placebo (19% for mepolizumab versus 1% for placebo; OR 19.7; 95% CI: 2.3, 167.9).(1)

The time to first relapse (defined as worsening related to vasculitis, asthma, or sinonasal symptoms requiring an increase in dose of corticosteroids or immunosuppressive therapy or hospitalization) was significantly longer for subjects receiving mepolizumab compared with placebo with a hazard ratio of 0.32 (95% CI: 0.21, 0.5). Additionally, subjects receiving mepolizumab had a reduction in rate of relapse compared with subjects receiving placebo (rate ratio 0.50; 95% CI: 0.36, 0.70 for mepolizumab compared with placebo). The incidence and number of relapse types (vasculitis, asthma, sino-nasal) were numerically lower with mepolizumab compared with placebo.(1)

Subjects receiving mepolizumab had a significantly greater reduction in average daily OCS dose compared with subjects receiving placebo during Weeks 48 to 52.(1)

HES

Nucala

A total of 108 adult and adolescent patients aged 12 years and older with HES for at least 6 months were evaluated in a randomized, double-blind, placebo-controlled, multicenter, 32-week trial (NCT #02836496). Patients with non-hematologic secondary HES (e.g., drug hypersensitivity, parasitic helminth infection, HIV infection, non-hematologic malignancy) or FIP1L1-PDGFRa kinase-positive HES were excluded from the trial. Patients received 300 mg of Nucala or placebo subcutaneously once every 4 weeks while continuing their stable HES therapy. Patients entering the trial had experienced at least 2 HES flares within the past 12 months and a blood eosinophil count of 1,000 cells/microliter or higher during screening. Historical HES flares for the trial entry criteria were defined as HES-related worsening of clinical symptoms or blood eosinophil counts requiring an escalation in therapy. Patients must have been on stable background HES therapy for a minimum of 4 weeks prior to randomization; existing HES therapy was maintained throughout the treatment period unless there was symptom worsening that required a dose increase. HES therapy could include chronic or episodic oral corticosteroids (OCS), immunosuppressive, and/or cytotoxic therapy.(1,10)

The efficacy of Nucala in HES was established based upon the proportion of patients who experienced a HES flare during the 32-week treatment period. A HES flare was defined as worsening of clinical signs and symptoms of HES or increasing eosinophils (on at least 2 occasions), resulting in the need to increase OCS or increase/add cytotoxic or immunosuppressive HES therapy. Over the 32-week treatment period, the incidence of HES flare over the treatment period was 56% for the placebo group and 28% for the group treated with Nucala (50% reduction).(1,10)

CRSwNP

Nucala

A randomized, double-blind, multicenter, placebo-controlled 52-week trial (NCT03085797) evaluated Nucala in patients with CRSwNP. The trial inclusion requirements included adult patients on background intranasal corticosteroids (INCS), with recurrent and symptomatic CRSwNP despite at least 1 surgery for the removal of nasal polyps within the previous 10 years. A total of 407 subjects were randomized to receive either 100 mg Nucala (N=206) or placebo (N=201) every 4 weeks for 52 weeks (13 doses). All study participants received mometasone furoate 400 mcg (intolerant participants received 200mcg) daily along with Nucala or placebo. Participants were not required to have sinus CT scans, but were required to have endoscopic confirmation of diagnosis.(1)

The co-primary efficacy endpoints were change from baseline to Week 52 in total endoscopic nasal polyps score (NPS; 0-8 scale) as graded by independent blinded assessors and change from baseline in nasal visual analog scale (VAS; 0-10 scale) during weeks 49 to 52.(1)

Statistically significant efficacy was observed regarding improvement (decrease) in bilateral endoscopic NPS score at week 52, and nasal obstruction VAS score from weeks 49 to 52. Total endoscopic NPS significantly improved at week 52 from baseline with mepolizumab versus placebo (adjusted difference in medians -0.73, 95% CI -1.11 to -0.34; p less than 0.001) and nasal obstruction VAS score during weeks 49–52 also significantly improved (-3.14, -4.09 to -2.18; p less than 0.001).(1)

Treatment with Nucala resulted in significant reduction of systemic corticosteroid use and need for sinonasal surgery versus placebo. The proportion of subjects who required surgery was reduced by 57% (HR of 0.43; 95% CI: 0.25, 0.76). Treatment

	with Nucala also significantly reduced the need for systemic steroids for nasal polyps versus placebo.(1)
Safety	 Fasenra (benralizumab) is contraindicated in those with known hypersensitivity to benralizumab or excipients.(2) Nucala (mepolizumab) is contraindicated in patients with history of hypersensitivity to mepolizumab or excipients in the formulation.(1) Benralizumab and mepolizumab have not been studied for use in combination with Xolair (omalizumab).

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POLICY AGENT SUMMARY PRIOR AUTHORIZATION

Target Brand Agent(s)	Target Generic Agent(s)	Strength	Targeted MSC	Available MSC	Final Age Limit	Preferred Status
Fasenra pen	benralizumab subcutaneous soln auto- injector	30 MG/ML	M ; N ; O ; Y	N		
Nucala	mepolizumab subcutaneous solution auto-injector	100 MG/ML	M ; N ; O ; Y	N		
Nucala	mepolizumab subcutaneous solution pref syringe	100 MG/ML ; 40 MG/0.4ML	M ; N ; O ; Y	N		

POLICY AGENT SUMMARY QUANTITY LIMIT

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strengt h	QL Amount	Dose Form	Day Supply	Duratio n	Addtl QL Info	Allowed Exceptions	Targete d NDCs When Exclusi ons Exist
Fasenra pen	Benralizumab Subcutaneous Soln Auto-injector 30 MG/ML	30 MG/ML	1	Pen	56	DAYS			
Nucala	Mepolizumab Subcutaneous Solution Auto- injector 100 MG/ML	100 MG/ML	3	Syringes	28	DAYS		Severe eosinophilic asthma: 1 syringe/28 days	
Nucala	Mepolizumab Subcutaneous Solution Pref Syringe	40 MG/0.4 ML	1	Syringe	28	DAYS			
Nucala	Mepolizumab Subcutaneous Solution Pref Syringe 100 MG/ML	100 MG/ML	3	Syringes	28	DAYS		Severe eosinophilic asthma: 1 syringe/28 days	

CLIENT SUMMARY – PRIOR AUTHORIZATION

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Client Formulary
Fasenra pen	benralizumab subcutaneous soln auto- injector	30 MG/ML	FlexRx Closed ; FlexRx Open ; FocusRx ; GenRx Closed ; GenRx Open ; Health Insurance Marketplace/BasicRx ; KeyRx

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Client Formulary
Nucala	mepolizumab subcutaneous solution auto-injector	100 MG/ML	FlexRx Closed ; FlexRx Open ; FocusRx ; GenRx Closed ; GenRx Open ; Health Insurance Marketplace/BasicRx ; KeyRx
Nucala	mepolizumab subcutaneous solution pref syringe	100 MG/ML ; 40 MG/0.4ML	FlexRx Closed ; FlexRx Open ; FocusRx ; GenRx Closed ; GenRx Open ; Health Insurance Marketplace/BasicRx ; KeyRx

CLIENT SUMMARY - QUANTITY LIMITS

Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	Client Formulary
Fasenra pen	Benralizumab Subcutaneous Soln Auto- injector 30 MG/ML	30 MG/ML	FlexRx Closed ; FlexRx Open ; FocusRx ; GenRx Closed ; GenRx Open ; Health Insurance Marketplace/BasicRx ; KeyRx
Nucala	Mepolizumab Subcutaneous Solution Auto-injector 100 MG/ML	100 MG/ML	FlexRx Closed ; FlexRx Open ; FocusRx ; GenRx Closed ; GenRx Open ; Health Insurance Marketplace/BasicRx ; KeyRx
Nucala	Mepolizumab Subcutaneous Solution Pref Syringe	40 MG/0.4ML	FlexRx Closed ; FlexRx Open ; FocusRx ; GenRx Closed ; GenRx Open ; Health Insurance Marketplace/BasicRx ; KeyRx
Nucala	Mepolizumab Subcutaneous Solution Pref Syringe 100 MG/ML	100 MG/ML	FlexRx Closed ; FlexRx Open ; FocusRx ; GenRx Closed ; GenRx Open ; Health Insurance Marketplace/BasicRx ; KeyRx

PRIOR AUTHORIZATION CLINICAL CRITERIA FOR APPROVAL

Module	Clinical Criteria for Approval				
	Initial Evaluation				
	Target Agent(s) will be approved when ALL of the following are met:				
	 ONE of the following: A. The requested agent is eligible for continuation of therapy AND ONE of the following: 				
	Agents Eligible for Continuation of Therapy				
	No Target Agents are Eligible for Continuation of Therapy				
	 Information has been provided that indicates the patient has been treated with the requested agent (starting on samples is not approvable) within the past 90 days OR The prescriber states the patient has been treated with requested agent (starting on samples is not approvable) within the past 90 days AND is at risk if therapy is changed OR B. The patient has a diagnosis of severe eosinophilic asthma and BOTH of the following: The patient's diagnosis has been confirmed by ONE of the following: 				

Module	Clinical Criteria for Approval
	A. The patient has a baseline (prior to therapy with the requested agent) blood eosinophilic count of 150 cells/microliter or higher while on high-dose inhaled corticosteroids or daily oral corticosteroids OR
	 B. The patient has a fraction of exhaled nitric oxide (FeNO) of 20 parts per billion or higher while on high-dose inhaled corticosteroids or daily oral corticosteroids OR
	 C. The patient has sputum eosinophils 2% or higher while on high- dose inhaled corticosteroids or daily oral corticosteroids AND
	 The patient has a history of uncontrolled asthma while on asthma control therapy as demonstrated by ONE of the following:
	 A. Frequent severe asthma exacerbations requiring two or more courses of systemic corticosteroids (steroid burst) within the past 12 months OR
	B. Serious asthma exacerbations requiring hospitalization, mechanical ventilation, or visit to the emergency room or urgent care within the past 12 months OR
	 C. Controlled asthma that worsens when the doses of inhaled and/or systemic corticosteroids are tapered OR
	D. The patient has baseline (prior to therapy with the requested agent) Forced Expiratory Volume (FEV1) that is less than 80% of predicted OR
	C. The patient has a diagnosis of eosinophilic granulomatosis with polyangiitis (EGPA) and ALL of the following:
	1. The requested agent is Nucala AND
	2. The patient has had a diagnosis of EGPA for at least 6 months with a
	history of relapsing or refractory disease AND
	3. The patient's diagnosis of EGPA was confirmed by ONE of the following:
	1. Asthma (history of wheezing or diffuse high-pitched rales
	on expiration)
	2. Eosinophilia (greater than 10% eosinophils on white blood cell differential count)
	 Mononeuropathy (including multiplex), multiple mononeuropathies, or polyneuropathy attributed to a systemic vasculitis
	4. Migratory or transient pulmonary infiltrates detected radiographically
	5. Paranasal sinus abnormality
	6. Biopsy containing a blood vessel showing the accumulation of eosinophils in extravascular areas OR
	B. The patient meets ALL of the following:
	1. Medical history of asthma AND
	2. Peak peripheral blood eosinophilia greater than 1500 cells/microliter AND
	3. Systemic vasculitis involving two or more extra-
	pulmonary organs AND
	A. The patient is currently on maximally tolerated oral corticosteroid therapy OR
	 B. The patient has an intolerance or hypersensitivity to oral corticosteroid therapy OR
	C. The patient has an FDA labeled contraindication to ALL oral corticosteroids AND
	A. The patient has tried and had an inadequate response to ONE
	non-corticosteroid immunosuppressant (e.g., azathioprine, cyclophosphamide, methotrexate, mycophenolate mofetil, rituvimab) OP
	B. The patient has an intolerance or hypersensitivity to ONE non- corticosteroid immunosuppressant OR

Module	Clinical Criteria for Approval
	C. The patient has an FDA labeled contraindication to ALL of the
	following immunosuppressants
	1. Azathioprine
	2. Cyclopnospnamide
	4. Mycophenolate mofetil OR
	D. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	1. A statement by the prescriber that the patient is currently
	taking the requested agent AND
	receiving a positive therapeutic outcome on requested
	agent AND
	3. The prescriber states that a change in therapy is expected
	to be ineffective or cause harm OR
	corticosteroid immunosuppressants cannot be used due to a
	documented medical condition or comorbid condition that is likely
	to cause an adverse reaction, decrease ability of the patient to
	achieve or maintain reasonable functional ability in performing
	daily activities of cause physical or mental narm OR The national has a diagnosis of hypereosinophilic syndrome (HES) and ALL of the
	following:
	1. The requested agent is Nucala AND
	2. BOTH of the following:
	A. The patient has had a diagnosis of HES for at least 6 months AND B. The patient has a history of at least 2 HES flares within the past
	12 months (i.e., worsening of clinical symptoms and/or blood
	eosinophil counts requiring an escalation in therapy) AND
	3. The patient's diagnosis of HES was confirmed by BOTH of the following:
	A. ONE of the following:
	areater than 1000 cells/microliter OR
	2. The patient has a percentage of eosinophils in bone
	marrow section exceeding 20% of all nucleated cells OR
	3. The patient has marked deposition of eosinophil granule
	4 The patient has tissue infiltration by eosinophils that is
	extensive in the opinion of a pathologist AND
	B. ALL of the following:
	1. Secondary (reactive, non-hematologic) causes of
	eosinophilia have been excluded (e.g., infection,
	metabolic [e.g., adrenal insufficiency], solid
	tumor/lymphoma) AND
	2. There has been evaluation of hypereosinophilia-related
	organ involvement (e.g., fibrosis of lung, heart, digestive
	cutaneous ervthema, edema/angioedema, ulceration.
	pruritis, or eczema; peripheral or central neuropathy with
	chronic or recurrent neurologic deficit; other organ
	system involvement such as liver, pancreas, kidney) AND
	disease OR
	E. The patient has a diagnosis of chronic rhinosinusitis with nasal polyposis
	(CRSwNP) AND ALL of the following:
	 The requested agent is Nucala AND The national bas at least TWO of the following symptoms consistent with
	chronic rhinosinusitis (CRS):
	A. Nasal discharge (rhinorrhea or post-nasal drainage)
	B. Nasal obstruction or congestion
	C. Loss or decreased sense of smell (hyposmia)

Module	Clinical Criteria for Approval
	D. Facial pressure or pain AND
	3. The patient has had symptoms consistent with chronic rhinosinusitis
	(CRS) for at least 12 consecutive weeks AND
	4. There is information indicating the patient's diagnosis was confirmed by
	ONE of the following:
	A. Anterior rhinoscopy or endoscopy OR
	B. Computed tomography (CT) of the sinuses AND
	5. ONE of the following:
	A. ONE of the following:
	1. The patient had an inadequate response to sinonasal
	surgery UK
	2. The patient is NOT a candidate for smortasal surgery UK
	B. ONE of the following. 1. The nation has tried and had an inadequate response to
	oral systemic corticosteroids OP
	The national has an intolerance or hypersensitivity to
	therapy with oral systemic corticosteroids OR
	3. The patient has an FDA labeled contraindication to AL
	oral systemic corticosteroids AND
	6. ONE of the following:
	A. The patient has tried and had an inadequate response to
	intranasal corticosteroids (e.g., fluticasone, Sinuva) OR
	B. The patient has an intolerance or hypersensitivity to therapy with
	intranasal corticosteroids (e.g., fluticasone, Sinuva) OR
	c. The patient has an FDA labeled contraindication to ALL intranasal
	corticosteroids OR
	F. The patient has another FDA approved indication for the requested agent and
	Foule of duministration OR
	requested agent and route of administration AND
	2. If the patient has a diagnosis of severe eosinophilic asthma, ALL of the following:
	A. ONE of the following:
	1. The patient is NOT currently being treated with the requested agent AND
	is currently treated with a maximally tolerated inhaled corticosteroid for
	at least 3 months OR
	2. The patient is currently being treated with the requested agent AND ONE
	of the following:
	A. Is currently treated with an innaled corticosteroid that is
	adequately dosed to control symptoms UK
	corticosteroid OP
	3 The patient has an intolerance or hypersensitivity to inhaled corticosteroid
	therapy OR
	4. The patient has an FDA labeled contraindication to ALL inhaled
	corticosteroids AND
	B. ONE of the following:
	1. The patient is currently being treated with ONE of the following:
	A. A long-acting beta-2 agonist (LABA) OR
	B. Long-acting muscarinic antagonist (LAMA) OR
	C. A leukotriene receptor antagonist (LTRA) OR
	2. The patient has an intolerance or hypersensitivity to therany with long-
	acting beta-2 agonists (LABA), long-acting muscarinic antagonists
	(LAMA), leukotriene receptor antagonists (LTRA) or theophylline OR
	3. The patient has an FDA labeled contraindication to ALL long-acting beta-2
	agonists (LABA) AND long-acting muscarinic antagonists (LAMA) AND
	C. The patient will continue asthma control therapy (e.g., ICS, ICS/LABA, LTRA,
	LAMA, theophylline) in combination with the requested agent AND
	3. If the patient has a diagnosis of hypereosinophilic syndrome (HES), ALL of the following:
	A. UNE of the following:
	1. The patient is currently being treated with maximally tolerated oral corticosteroid (OCS) OP
1	

Module	Clinical Criteria for Approval
	2. The patient has an intolerance or hypersensitivity to oral corticosteroid
	(OCS) therapy OR
	3. The patient has an FDA labeled contraindication to ALL oral corticosteroids OR
	4. The patient is currently being treated with the requested agent as
	A A statement by the prescriber that the patient is currently taking
	the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutic outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be
	The prescriber has provided documentation that ALL oral
	corticosteroids cannot be used due to a documented medical condition or
	comorbid condition that is likely to cause an adverse reaction, decrease
	ability of the patient to achieve or maintain reasonable functional ability
	in performing daily activities or cause physical or mental harm AND
	B. ONE of the following:
	A. Hydroxyurea OR
	B. Interferon-a OR
	C. Another immunosuppressive agent (e.g., azathioprine,
	cyclosporine, methotrexate, tacrolimus) OR
	2. The patient has an incolerance of hypersensitivity to therapy with hydroxyurea, interferon-q, or immunosuppressive agents (e.g.,
	azathioprine, cyclosporine, methotrexate, tacrolimus) OR
	3. The patient has an FDA labeled contraindication to hydroxyurea,
	interferon-a, and ALL immunosuppressive agents (e.g., azathioprine,
	cyclosporine, methotrexate, tacrolimus) OR
	indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking
	the requested agent AND
	B. A statement by the prescriber that the patient is currently
	C The prescriber states that a change in therapy is expected to be
	ineffective or cause harm OR
	5. The prescriber has provided documentation that hydroxyurea, interferon-
	a, and ALL immunosuppressive agents (e.g., azathioprine, cyclosporine,
	methotrexate, tacrolimus) cannot be used due to a documented medical
	reaction decrease ability of the nation to achieve or maintain reasonable
	functional ability in performing daily activities or cause physical or mental
	harm AND
	C. The patient will continue existing HES therapy (e.g., OCS, hydroxyurea,
	Interferon-d, Immunosuppressants) in combination with the requested agent AND
	BOTH of the following:
	A. The patient is currently treated with standard nasal polyp maintenance therapy
	(e.g., nasal saline irrigation, intranasal corticosteroids) AND
	B. The patient will continue standard nasal polyp maintenance therapy (e.g., nasal
	saline irrigation, intranasal corticosteroids) in combination with the requested agent AND
	5. If the patient has an FDA approved indication, then ONE of the following:
	A. The patient's age is within FDA labeling for the requested indication for the
	requested agent OR
	B. The prescriber has provided information in support of using the requested agent for the patient's age for the requested indication AND .
	6. The prescriber is a specialist in the area of the nation's diagnosis (e.g., allergist
	immunologist, otolaryngologist, pulmonologist) or the prescriber has consulted with a
	specialist in the area of the patient's diagnosis AND
	7. ONE of the following (Please refer to "Agents NOT to be used Concomitantly" table):

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Module	Clinical Criteria for Approval
	D. Dose of maintenance corticosteroid therapy and/or
	immunosuppressant therapy was not increased AND
	A. The patient is currently treated and is compliant with
	maintenance therapy with oral corticosteroids OR
	B. The patient has an intolerance or hypersensitivity to oral
	corticosteroid therapy OR
	corticosteroids OR
	D. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	1. A statement by the prescriber that the patient is currently
	2. A statement by the prescriber that the patient is currently
	receiving a positive therapeutic outcome on requested
	agent AND
	 The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	E. The prescriber has provided documentation that ALL oral
	conticosteroids cannot be used due to a documented medical
	reaction, decrease ability of the patient to achieve or maintain
	reasonable functional ability in performing daily activities or cause
	physical or mental harm OR
	C. The patient has a diagnosis of hypereosinophilic syndrome (HES) AND ALL of the following:
	1. The requested agent is Nucala AND
	2. The patient has had improvements or stabilization with the requested
	agent from baseline (prior to therapy with the requested agent) as
	A Decrease in incidence of HES flares OR
	B. Escalation of therapy (due to HES-related worsening of clinical
	symptoms or increased blood eosinophil counts) has not been
	required AND
	A. The patient is currently treated and is compliant with oral
	corticosteroid and/or other maintenance therapy (e.g.,
	hydroxyurea, interferon-a, azathioprine, cyclosporine,
	methotrexate, tacrolimus) OR
	oral corticosteroids or other maintenance agents (e.g.,
	hydroxyurea, interferon-a, azathioprine, cyclosporine,
	methotrexate, tacrolimus) OR
	C. The patient has an FDA labeled contraindication to ALL oral
	interferon-q, azathioprine, cyclosporine, methotrexate,
	tacrolimus) OR
	D. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	taking the requested agent AND
	2. A statement by the prescriber that the patient is currently
	agent AND
	3. The prescriber states that a change in therapy is expected
	to be ineffective or cause harm OR
	E. The prescriber has provided documentation that ALL oral
	interferon-q, azathionrine, cyclosporine, methotrexate
	tacrolimus) cannot be used due to a documented medical
	condition or comorbid condition that is likely to cause an adverse
	reaction, decrease ability of the patient to achieve or maintain

	Clinical Criteria for Approval
	reasonable functional ability in performing daily activities or cause
	physical or mental harm OR
	D. The patient has a diagnosis of chronic rhinosinusitis with nasal polyposis
	(CRSwNP) AND ALL of the following:
	1. The requested agent is Nucala AND
	2. The patient has had clinical benefit with the requested agent AND
	3. The patient will continue standard hasal polyp maintenance therapy (e.g.,
	nasal saline irrigation, intranasal corticosteroids) in combination with the
	requested agent OR
	E. The patient has another FDA approved indication for the requested agent and
	route of administration AND has had clinical benefit with the requested agent OK
	F. The patient has another indication that is supported in compendia for the
	requested agent AND
	The proceriber is a specialist in the area of the patient's diagnosis (e.g., allergist
	immunologist otologist nulmonologist) or the proscriber has consulted with a
	specialist in the area of the patient's diagnosis AND
	4 ONE of the following (Please refer to "Agents NOT to be used Concomitantly" table):
	The patient will NOT be using the requested agent in combination with another
	immunomodulatory agent (e.g. TNE inhibitors 1AK inhibitors 11-4 inhibitors) OR
	B The patient will be using the requested agent in combination with another
	immunomodulatory agent AND BOTH of the following:
	1. The prescribing information for the requested agent does NOT limit the
	use with another immunomodulatory agent AND
	2. The prescriber has provided information in support of combination
	therapy (submitted copy required, e.g., clinical trials, phase III studies,
	guidelines required) AND
	5. The patient does NOT have any FDA labeled contraindications to the requested agent
Co	ompendia Allowed: AHFS, DrugDex 1 or 2a level of evidence, or NCCN 1 or 2a recommended
us	e
Le	ength of Approval: 12 months
NC	JIE: If Quantity Limit applies, please refer to Quantity Limit Criteria.

<u>QUANTI</u>	TY LIMIT CLINICAL CRITERIA FOR APPROVAL
Module	Clinical Criteria for Approval
	Quantity Limit for the Target Agent(s) will be approved when ONE of the following is met:
	 The requested quantity (dose) does NOT exceed the program quantity limit OR ALL of the following: A. The requested quantity (dose) exceeds the program quantity limit AND B. The requested quantity (dose) does NOT exceed the maximum FDA labeled dose for the requested indication AND C. The requested quantity (dose) cannot be achieved with a lower quantity of a higher strength that does not exceed the program quantity limit
	Length of Approval: Initial: 6 months for severe eosinophilic asthma; 12 months for EGPA, HES, CRSwNP, and all other FDA approved indications; For Fasenra, approve loading dose for new starts and the maintenance dose for the remainder of the 6 months; Renewal: 12 months

CONTRAINDICATION AGENTS

Contraindicated as Concomitant Therapy

Agents NOT to be used Concomitantly

Contraindicated as Concomitant Therapy Abrilada (adalimumab-afzb) Actemra (tocilizumab) Adalimumab Adbry (tralokinumab-ldrm) Amjevita (adalimumab-atto) Arcalyst (rilonacept) Avsola (infliximab-axxq) Benlysta (belimumab) Bimzelx (bimekizumab-bkzx) Cibinqo (abrocitinib) Cimzia (certolizumab) Cinqair (reslizumab) Cosentyx (secukinumab) Cyltezo (adalimumab-adbm) Dupixent (dupilumab) Enbrel (etanercept) Entyvio (vedolizumab) Fasenra (benralizumab) Hadlima (adalimumab-bwwd) Hulio (adalimumab-fkjp) Humira (adalimumab) Hyrimoz (adalimumab-adaz) Idacio (adalimumab-aacf) Ilaris (canakinumab) Ilumya (tildrakizumab-asmn) Inflectra (infliximab-dyyb) Infliximab

Contra	vindicated as Concomitant Therany
Kevza	ara (sarilumab)
Kiner	et (anakinra)
Litful	o (ritlecitinib)
Nuca	a (mepolizumab)
Olum	iant (baricitinib)
Omvo	oh (mirikizumab-mrkz)
Opze	ura (ruxolitinib)
Oren	cia (abatacept)
Otezl	a (apremilast)
Remi	cade (infliximab)
Renfl	exis (infliximab-abda)
Riabr	i (rituximab-arrx)
Rinvo	q (upadacitinib)
Ritux	an (rituximab)
Ritux	an Hycela (rituximab/hyaluronidase human)
Ruxie	nce (rituximab-pvvr)
Siliq	(brodalumab)
Simp	oni (golimumab)
Simp	oni ARIA (golimumab)
Skyri	zi (risankizumab-rzaa)
Sotyk	tu (deucravacitinib)
Stela	ra (ustekinumab)
Taltz	(ixekizumab)
Tezsp	ire (tezepelumab-ekko)
Trem	fya (guselkumab)
Truxi	ma (rituximab-abbs)
Tysat	pri (natalizumab)

Contraindicated as Concomitant Therapy
Velsipity (etrasimod)
Wezlana (ustekinumab-auub)
Xeljanz (tofacitinib)
Xeljanz XR (tofacitinib extended release)
Xolair (omalizumab)
Yuflyma (adalimumab-aaty)
Yusimry (adalimumab-aqvh)
Zeposia (ozanimod)
Zymfentra (infliximab-dyyb)